WEST

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L4: Entry 6 of 9

File: USPT

Nov 16, 1993

US-PAT-NO: 5262325

DOCUMENT-IDENTIFIER: US 5262325 A

TITLE: Method for the enzymatic neutralization of heparin

DATE-ISSUED: November 16, 1993

INVENTOR-INFORMATION:

NAME CITY STATE ZIP CODE COUNTRY

Zimmermann; Joseph J. Elm Grove WI

Lewis; N. Tracey Brossard CAX
Heft; Robert A. Ville St. Laurent CAX

US-CL-CURRENT: 435/269; 435/13, 435/200

CLAIMS:

We claim:

1. A method to eliminate the physiological effects of heparin on blood components in a mixture of blood components and heparin comprising

contacting blood components containing heparin with a bacterial heparinase preparation,

wherein the $\underline{\text{heparinase}}$ preparation is free of an anticoagulant component prior to mixing of the heparinase preparation with the heparin,

wherein the anticoagulant component does not bind to a polysulfated resin at pH 7.0, conductivity between 3 and 12 mmhos, and the <u>heparinase</u> does bind to a polysulfated resin at pH 7.0, conductivity between 3 and 12 mmhos, and

wherein the <u>heparinase</u> preparation free of an anticoagulant component has an optimal activity at pH=6.5 to 7.0; salt concentration=0.1M, and 37.degree. C.

- 2. The method of claim 1, wherein the <u>heparinase</u> is purified from cultures of Flavobacterium heparinum chromatography using a polysulfated resin.
- 3. The method of claim 2 wherein the <u>heparinase</u> is eluted from the affinity resin by increasing the conductivity of the eluent such that the eluted <u>heparinase</u> fraction is free of a Flavobacterium heparinum component inhibiting coagulation.
- 4. The method of claim 1 wherein the blood contains low molecular weight heparinum.
- 5. The method of claim 1 wherein the <u>heparinase</u> formulation is prepared by lyophilizing 0.05 to 3.0 IU anticoagulant free <u>heparinase</u> in the presence of 0.5-1.0 mg ammonium sulfate/IU <u>heparinase</u> which has been placed within containers suitable for the collection or incubation of blood or plasma samples.
- 6. The method of claim 5 wherein the containers are selected from the group consisting of blood collection tubes, pipet tips, fluid transfer devices and syringes.

- 7. The method of claim 1 wherein the blood sample is incubated with the lyophilized heparinase formulation for between 1 and 3600 seconds at a temperature between 2.degree. and 41.degree. C. prior to the initiation of a hematological test.
- 8. The method of claim 7 wherein the test relies on the formation of a blood clot as its endpoint.
- 9. The method of claim 1 wherein the heparinase formulation is prepared by lyophilizing 0.05 to 300 IU anticoagulant free heparinase in the presence of 0.5-1.0 mg ammonium sulfate/IU heparinase which has been placed within a container further comprising resolubilizing the heparinase with a physiologically compatible solution.
- 10. The method of claim 1 further comprising injecting intravenously the dissolved heparinase formulation into a patient receiving heparin therapy for the purpose of neutralizing the anticoagulant properties of heparin in the bloodstream.
- 11. The method of claim 10 wherein the heparin is low molecular weight heparin.

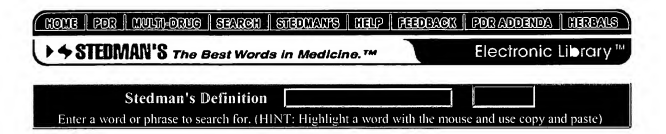
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vein (v?n) [TA]

A blood vessel carrying blood toward the heart; postnatally, all v. except the pulmonary carry dark unoxygenated blood. SYN: vena [TA]. [L. vena] accessory cephalic v. [TA] a variable v. that passes along the radial border of the forearm to join the cephalic v. near the elbow. SYN: vena cephalica accessoria [TA]. accessory hemiazygos v. [TA] formed by the union of the fourth to seventh left posterior intercostal v., passes along the side of the bodies of the fifth, sixth, and seventh thoracic vertebrae, then crosses the midline behind the aorta, esophagus, and thoracic duct, and empties into the azygos v., sometimes in common with the hemiazygos v.. SYN: vena hemiazygos accessoria [TA], vena azygos minor superior. accessory saphenous v. [TA] an occasional v. running in the thigh parallel to the great saphenous ν , which it joins just before the latter empties into the femoral ν . SYN: vena saphena accessoria [TA]. accessory vertebral v. [TA] a v. that accompanies the vertebral v. but passes through the foramen of the transverse process of the seventh cervical vertebra and opens independently into the brachiocephalic v.. SYN: vena vertebralis accessoria [TA]. accompanying v. SYN: vena comitans. accompanying v. of hypoglossal nerve SYN: vena comitans of hypoglossal nerve. anastomotic v. inferior anastomotic v., superior anastomotic v.. angular v. [TA] a short v. at the medial angle of the eye, formed by the supraorbital and supratrochlear v. and continuing as the facial v. SYN: vena angularis [TA] anonymous v. obsolete term for (left and right) brachiocephalic v.. anterior auricular v. [TA] one of several v. draining the auricle and acoustic meatus and emptying into the retromandibular v... SYN: vena auricularis anterior, vena preauricularis, anterior basal v. [TA] SYN: vena basalis anterior [TA], anterior basal branch of superior basal vein (of right and left inferior pulmonary veins)&star, ramus basalis anterior venae basalis superioris&star. anterior cardiac v. [TA] two or three small v. in the anterior wall of the right ventricle opening directly into the right atrium independently of the coronary sinus. SYN: venae cardiacae anteriores [TA]. anterior cerebral v. [TA] small v. that parallel the anterior cerebral artery and drain into the basal v.. SYN: venae anteriores cerebri [TA]. anterior ciliary v. [TA] several small v., anterior and posterior, coming from the ciliary body. SYN: venae ciliares anteriores [TA]. anterior circumflex humeral v. [TA] vein accompanying the artery of the same name, passing anterior to the surgical neck of the humerus to enter the axillary vein. SYN: vena circumflexa humeri anterior [TA] . anterior facial v. SYN: facial v. anterior intercostal v. [TA] tributaries to the musculophrenic or internal thoracic v. from the

1 of 22



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intravenous (I.V., i.v.)

Within a vein or veins. SYN: endovenous.

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intravascular

Within the blood vessels or lymphatics.

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WEST Search History

DATE: Wednesday, May 29, 2002

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DB=US	SPT,PGPB,JPAB,EPAB,DWPI; PLUR=YES; OP=OR		
L7	L6 near (intravascul\$9)	0	L7
L6	L5 not 14	3	L6
L5	heparinase\$ near administ\$8	7	L5
L4	L3 and administ\$6	9.	L4
L3	L2 and heparinase\$	24	L3
L2	(bennett)[IN] OR (cauchon)[IN] or (fink)[in] or (grouix)[in] or (hsia)[in] or (danagher)[in] or (zimmermann)[in]	18403	L2
L1	(bennett)[IN] OR (cauchon)[IN]	8486	L1

END OF SEARCH HISTORY

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NEWS 2
                                     frequency
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                                    TOXLIT no longer available
TRCTHERMO no longer available
                   Mar 22
Mar 22
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                    Mar 28
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                    Apr 02
Apr 08
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   NEWS 15 Apr 19
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                                     Federal Research in Progress (FEDRIP) now available
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AND CURRENT DISCOVER FILE IS DATED 05 FEBRUARY 2002
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                     3125 BENNET D?/AU OR CAUCHON E?/AU OR FINK D?/AU OR GROUIX B?/AU OR HSIA A?/AU OR DANAGHER P?/AU OR ZIMMERMANN J?/AU
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FIELD CODE - 'AND' OPERATOR ASSUMED 'L19 (P)
L5 0 L4 (P) (IV OR INTRAVASCULAR?)
                           0 L4 (P) (IV OR INTRAVASCULAR?)
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=> dis 14 1-7 ibib abs ANSWER 1 OF 7 MEDITINE ACCESSION NUMBER: MEDLINE PubMed ID: 8712450 DOCUMENT NUMBER: 96322828 Heparinase I (neutralase) reversal of systemic anticoaculation. anticoagulation.
Michelsen L G, Kikura M, Levy J H, Lee M K, Lee K C,
Zimmermann J J, Szlam F
Department of Anesthesiology, Emory University School of
Medicine, Atlanta, Georgia, USA.
ANESTHESIOLOGY, (1996 Aug) 85 (2) 339-46.
Journal code: 4SG; 1300217. ISSN: 0003-3022.
United States
JOURNAL Article. (TOURNAL ARTICE) AUTHOR: CORPORATE SOURCE: SOURCE: PUB. COUNTRY: Journal; Article; (JOURNAL ARTICLE) Abridged Index Medicus Journals; Priority Journals 199609 LANGUAGE: FILE SEGMENT: ENTRY MONTH: NY MONTH: 199609

Last Updated on STN: 19980206

Entered Medline: 19960912

BACKGROUND: Protamine causes multiple adverse reactions. Heparinase I, a specific enzyme that inactivates heparin, is a possible alternative to protamine. In this study, the authors examined the efficacy of heparinase I to reverse heparin-induced anticoagulation in vitro and compared heparinase I to protamine as an antagonist of heparin-induced anticoagulation in dogs. METHODS: In the in vitro study, blood was obtained from the extracorporeal circuits of 12 patients, and activated clotting times were determined after adding different concentrations of heparinase I. In the in vivo study, 24 anesthetized dogs received 300 units/kg heparin injected intravenously for 5 s, then 10 min later, 3.9 mg/kg protamine, 5-41 micrograms/kg heparinase I, or the vehicle (n = 4/group) were administered intravenously, and activated clotting times and hemodynamics were measured. RESULTS: In the in vitro study, heparin concentrations of 3.3 +/- 1.0 (mean +/- SD) units/ml (approximately 0.033 mg/ml; n = 12) were reversed in the blood of patients by heparinase I at concentrations > 0.490 microgram/ml. In the canine study, heparinase at all doses studied and protamine effectively reversed the anticoagulating effects of heparin within 10 min of administration. Protamine produced adverse hemodynamic effects, whereas heparinase or its vehicle produced no significant change in arterial pressure. CONCLUSION: Both heparinase I and protamine effectively reversed heparin anticoagulation. However, as opposed to protamine, heparinase I did not produce any significant hemodynamic changes when given as a bolus to dogs.

ANSWER 2 OF 7 MEDLINE Entered STN: 19960919 ENTRY DATE: 95030369 MEDLINE
95030369 PubMed ID: 7943773
In vitro reversal of heparin effect with heparinase: evaluation with whole blood prothrombin time and activated partial thromboplastin time in cardiac ACCESSION NUMBER: DOCUMENT NUMBER: TITLE: and activated partial thromboplastin time in cardiac surgical patients.
Despotis G J; Summerfield A L; Joist J H; Goodnough L T; Santoro S A; Zimmermann J J; Lappas D G
Department of Anesthesiology, Washington University School of Medicine, St. Louis, MO 63110.
ANESTHESIA AND ANALGESIA, (1994 Oct) 79 (4) 670-4.
Journal code: 4R8, 1310650. ISSN: 0003-2999.
United States AUTHOR: CORPORATE SOURCE: SOURCE . PUB. COUNTRY: United States (CLINICAL TRIAL) Journal; Article; (JOURNAL ARTICLE) LANGUAGE: English FILE SEGMENT: Abridged Index Medicus Journals; Priority Journals SEGMENT: Abridged Index Medicus Journals; Priority Journals Y MONTH: 199411
Y DATE: Entered STN: 19941222
Last Updated on STN: 19980206
Entered Medline: 19941110
This study was designed to evaluate the potential in vitro use of heparinase to eliminate functionally active heparin prior to performing whole blood (WB) prothrombin time (PT) and activated partial thromboplastin time (APTT) assays. A total of 250 U/kg of heparin for cardiopulmonary bypass (CPB) was administered to 30 cardiac surgical patients in three consecutive, divided doses (20, 80, and 150 U/kg) at 15-min intervals. Blood specimens were obtained prior to heparin administration (baseline) and 10 min after each heparin dose. After collection, blood specimens were fractionated into three aliquots of which the first was used for determination of heparin concentration. After gentle mixing, WB PT and APTT measurements were performed for heparinase (Aliquot 2)- and nonheparinase (Aliquot 3)-treated blood. With consecutive heparin doses of 20 and 80 U/kg, WB PT increased from a baseline of 12.3 +/-0.1 s to 13.3 +/-0.2 and 18.5 +/-1.3 s, while WB APTT increased from a baseline of 28.3 +/-1.1 s to 89.5 +/-5.4 after the initial heparin dose (20 U/kg). When compared to baseline (no heparin) results, small, progressive increases in heparinase-treated WB PT (0.7 +/-0.1, 1.5 +/-0.1, 2.1 +/-0.1 s) and APTT (2.3 +/-0.3, 5.7 +/-0.4, 9.5 +/-0.5 s) were seen with increasing heparin concentration (0.23, 1.58, and 3.95 U/mL, respectively). Heparinase was highly effective in eliminating the anticoagulant effects of even large amounts of heparin in plasma from cardiac surgical patients. (ABSTRACT TRUNCATED AT 250 WORDS) ENTRY MONTH: 199411 ANSWER 3 OF 7 CAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 1997:303407 CAPLUS DOCUMENT NUMBER: 126:272357 Use of heparinases to decrease inflammatory responses TITLE: Bennett, D. Clark; Cauchon, Elizabeth; Fink, Dominique; Grouix, Brigette; INVENTOR(S): Hsia, Ariane; Danagher, Pamela; Zimmerman, Joseph Ibex Technologies Inc., Can.; Bennett, D. Clark; PATENT ASSIGNEE(S):

PATENT ASSIGNEE(S): Ibex Technologies Inc., Can.; Bennett, D. Clark; Cauchon, Elizabeth, Fink, Dominique; Grouix, Brigette; Hsia, Ariane; Danagher, Pamela; Zimmerman, Joseph PCT Int. Appl., 73 pp. CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

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96-US15593 19960927
                  WO 9711684
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                              9711684 A1 19970403 W0 7996-US15993 19960927
W: AU, BR, CA, CM, CZ, HU, IL, JP, KR, MX, NO, NZ, SG, US, VN, AM,
AZ, BY, KG, KZ, MD, RU, TJ, TM
RW: AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE
2233343 AA 19970403 CA 1996-2233343 19960927
9673791 A1 19970417 AU 1996-73791 19960927
703394 B2 19990325
B22401 h1 19980315 FP 1996-936052 19960927
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                                                                                                                                                        EP 1996-936052
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                              AI 19780715 EP 1996-936052 19960927
R: AT, BE, CH, DE, DK, ES, PR, GB, GR, IT, LI, LU, NL, SE, MC, PT,
IE, FI
            JP 11512721 T2 19991102 JP 1996-513723 19960927
US 2001006635 A1 20010705 US 1996-722659 19960927
US 1995-4622P P 19950929
WO 1996-US15593 W 19960927
Heparinase enzymes can be used as a medical treatment to reduce localized inflammatory responses. Treatment of activated endothelium with heparinase inhibits leukocyte rolling, adhesion and extravasation. Most of the heparin and heparan sulfate on endothelial cell surfaces and in basement membranes is degraded by exposure to heparinase. In addn., immobilized chemokines, which are attached to heparin
/heparan sulfate on activated endothelium are solubilized by heparinase digestion. Heparinase can be infused into the vascular system to inhibit accumulation of leukocytes in inflamed tissue and decrease damage resulting from localized inflammations. Targeting of heparinase to activated endothelium can be accomplished through localized administration and/or use of genetically engineered heparinase contg. endothelium ligand-binding domains.

ANSWER 4 OF 7 CAPLUS COPVELGUE 2002 157
PRIORITY APPLN. INFO.:
                 ANSWER 4 OF 7 CAPLUS COPYRIGHT 2002 ACS
ACCESSION NUMBER:
DOCUMENT NUMBER:
                                                                                        1996:237498 CAPLUS
124:250963
                                                                                        124:250963
Glycosaminoglycan-degrading enzymes for modulation of wound healing processes
Zimmermann, Joseph; Vlodavsky, Israel;
Bennett, D. Clark; Danagher, Pamela;
Broughton, Richard
Ibex Technologies R and D, Inc., USA
PCT Int. Appl., 83 pp.
CODEN: PIXXD2
TITLE:
INVENTOR (S):
PATENT ASSIGNEE(S):
SOURCE:
                                                                                          English
 LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
                  PATENT NO.
                                                                             KIND DATE
                                                                                                                                                        APPLICATION NO. DATE
                  WO 9601648
                              A1
                                                                                                  19960125
                                                                                                                                                         WO 1995-US8608 19950707
                  US 5997863
                  CA 2194370
                 AU 9530949 Al 19960209
AU 707007 B2 19990701
EP 769961 Al 19970502 EP 1995-926645 19950707
R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, MC, NL, PT, SE
JP 10506609 T2 19980630 JP 1995-50443 19950707
RITY APPLN. INFO:
US 1994-273109 19940708
W0 1995-US8608 19950707
W0 1995-US8608 19950707
                   AU 9530949
                                                                                                                                                         AU 1995-30949
 PRIORITY APPLN. INFO.:
              OS 1994-273109 19940/08 WO 1995-US8608 19950707
Glycosaminoglycan-degrading enzymes, including heparinases 1, 2, and 3, as well as chondroitinases AC and B from the Gram neg. bacterial Flavobacterium heparinum, can be used either sep. or in combination to manipulate cell proliferation. In one embodiment, heparinases are administered to degrade heparan sulfate components of the extracellular matrix, thereby allowing the heparin-binding growth factors which are stored in the extracellular matrix to migrate to adjacent cells. The mobility of chemoattractant agents, growth factors and cells also can be increased by treating tissues with glycosaminoglycan-degrading enzymes, both chondroitinases and heparinases. The enzymic removal of chondroitin sulfates from cell surfaces effectively increases the availability of growth factor receptors on the cell's surface. Selectively removing heparan sulfate from cell surfaces while leaving the extracellular matrix intact, conversely, inhibits cell proliferation by down-regulating the cell's response to growth factors. This is achieved by targeting heparin- or heparan sulfate-degrading activities to the cell surface. Targeting the heparin-degrading activities to
                  or by phys. con administration.
L4 ANSWER 5 OF 7 CAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 1993:11727 CAPLUS
 DOCUMENT NUMBER:
                                                                                          118:11727
                                                                                          Heparinase from Plavobacterium heparinum for heparin neutralization
Zimmermann, Joseph J.; Lewis, N. Tracey;
 INVENTOR (S):
                                                                                          Heft, Robert A.
                                                                                          Hert, Robert A.

Ibex Technologies, Inc., Can.

PCT Int. Appl., 28 pp.

CODEN: PIXXD2

Patent
 PATENT ASSIGNEE(S):
 SOURCE:
 LANGUAGE:
                                                                                          English
 PAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
                   PATENT NO.
                                                                               KIND DATE
                                                                                                                                                          APPLICATION NO. DATE
                   WO 9217203
                                                                                  A1 19921015
                                                                                                                                                           WO 1992-US2724 19920403
                 W: AU, CA, JP
RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LU, MC, NL, SE
US 5262325
A 19931116
US 1991-680330
19910404
AU 9217713
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EP 537325
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                   R: AT, BE, CH, DE, DK, ES, FR, GB, JP 05507297 T2 19931021 J
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US 1993-153134

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PRIORITY APPLN. INFO.:
                                         US 1991
                                                              19910404
                                         WO 1992-US2724
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WO 1992-USZ724 19920403
A heparinase derived from F. heparinum eliminates heparin
interference of normal blood function. The heparinase is free of a
component that inhibits coagulation. The heparinase is useful in vitro to
eliminate the interference in hematol. assays due to the presence of
heparin and for the in vivo neutralization of heparin
during surgical procedures. It achieves neutralization faster and more
completely than previous compns. and is stable for a long time. When
rabbits were administered 250 IU heparin/kg, the
activated clotting time (ACT) was increased from 180 to >999 s. When 2.5
IU heparinase/kg was injected over the next h, the ACT returned to normal.
Heparinase was purified by affinity chromatog. and was eluted with a
concd. eluent to elute a fraction free of the coagulation-inhibition
component. ANSWER 6 OF 7 CAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 1989:13443 CAPLUS Incompatibility of dacarbazine and heparin sodium DOCUMENT NUMBER: AUTHOR(S): CORPORATE SOURCE: Schroeder, Frank; Zimmermann, Jutta Zentralapotheke, Zentralkrankenhaus., Bremen, 2800/1, Fed. Rep. Ger. Pharm. Ztg. (1988), 133(36), 24, 26 CODEN: PHZIAP; ISSN: 0031-7136 SOURCE: DOCUMENT TYPE: Journal LANGUAGE: German The use of dacarbazine in oncol. studies leads to massive pain reactions and to decrease the pain heparin Na salt is usually administered i.v. The direct contact of both these drugs in an administered i.v. The direct contact of both these drugs in an infusion system leads to the formation of turbidity of the solns. in catheters. As a temporary soln. to this problem, the administration of dacarbazine in the form of short infusion while avoiding the simultaneous heparin administration is BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC. 2000:292364 BIOSIS ANSWER 7 OF 7 ACCESSION NUMBER: DOCUMENT NUMBER: PREV200000292364 Attenuation of wound healing processes.

Zimmermann, Joseph (1); Vlodavsky, Israel;
Bennett, Clark; Danagher, Pamel; Broughton, TITLE: AUTHOR (S) Richard CORPORATE SOURCE: (1) Montreal Canada ASSIGNEE: Ibex Technologies R and D, Inc., Montreal, Canada US 5997863 December 07, 1999
Official Gazette of the United States Patent and Trademark Office Patents, (Dec. 7, 1999) Vol. 1229, No. 1, pp. No pagination. e-file. PATENT INFORMATION: SOURCE: ISSN: 0098-1133. MENT TYPE: Patent
UAGE: English
Glycosaminoglycans, including heparinases 1, 2 and 3 as well as chondroitinases AC and B from the Gram negative bacteria Flavobacterium heparinum, can be used either separately or in combination to manipulate cell proliferation. In one embodiment, heparinases are administered to degrade heparan sulfate components of the extracellular matrix, thereby allowing the heparin binding growth factors which are stored in the extracellular matrix to migrate to adjacent cells. The mobility of chemoattractant agents, growth factors and cells also can be increased by treating tissues with glycosaminoglycan degrading enzymes, both chondroitinases and heparinases. The enzymatic removal of chondroitin sulfates from cell surfaces effectively increases the availability of growth factor receptors on the cell's surface. Selectively removing heparan sulfate from cell surfaces while leaving the extracellular matrix intact, conversely, inhibits cell proliferation by down regulating the cell's response to growth factors. This is achieved by targeting heparin or heparan sulfate degrading activity can be achieved by genetically engineering a ligand binding functionality into the heparinase proteins, or by physically controlling the localized enzyme concentration through the method of administration. DOCUMENT TYPE: Patent LANGUAGE: (FILE 'HOME' ENTERED AT 09:28:02 ON 29 MAY 2002) FILE 'MEDLINE, CAPLUS, EMBASE, BIOSIS' ENTERED AT 09:28:14 ON 29 MAY 2002 3125 S BENNET D?/AU OR CAUCHON E?/AU OR FINK D?/AU OR GROUIX B?/AU O 43 S L1 AND HEPARIN OR HEPARINSE L1 L2 21 DUP REM L2 (22 DUPLICATES REMOVED)
7 S L3 (P) ADMINIST?
0 S L4 (P) (IV OR INTRAVASCULAR?) L3 L5 s 11 and (heparinase or heparin)
59 L1 AND (HEPARINASE OR HEPARIN) => dup rem 16
PROCESSING COMPLETED FOR L6
L7 29 DUP REM L6 (30 DUPLICATES REMOVED) => S 17 (P) administ?
PROXIMITY OPERATOR LEVEL NOT CONSISTENT WITH
PIELD CODE - 'AND' OPERATOR ASSUMED 'L32 (P) ADMINIST?'
PROXIMITY OPERATOR LEVEL NOT CONSISTENT WITH
PIELD CODE - 'AND' OPERATOR ASSUMED 'L34 (P) ADMINIST?'
PROXIMITY OPERATOR LEVEL NOT CONSISTENT WITH
PIELD CODE - 'AND' OPERATOR ASSUMED 'L36 (P) ADMINIST?'
PROXIMITY OPERATOR LEVEL NOT CONSISTENT WITH
PIELD CODE - 'AND' OPERATOR ASSUMED 'L38 (P) ADMINIST?'
L8 7 L7 (P) ADMINIST? => s 18 not 14 L9 0 L8 NOT L4

s (heparinase or heparinase?)
3653 (HEPARINASE OR HEPARINASE?) => s 110 (10N) (administ? UNMATCHED LEFT PARENTHESIS '10A) (ADMINIST?' The number of right parentheses in a query must be equal to the number of left parentheses. s l10 (10N) (administ?) 1 41 L10 (10N) (ADMINIST?) => s 111 (P) (iv or intravascul?) 0 L11 (P) (IV OR INTRAVASCUL?)

=> dup rem 111

PROCESSING COMPLETED FOR L11
L13 18 DUP REM L11 (23 DUPLICATES REMOVED)

=> s 113 not 14 L14 14 L13 NOT L4

=> dis l14 1-14 ibib abs kwic

ANSWER 1 OF 14

ACCESSION NUMBER: DOCUMENT NUMBER:

TITLE:

MEDLINE
201662061 MEDLINE
201662061 MEDLINE
21583162 PubMed ID: 11726421
A dose-determining trial of heparinase-I (Neutralase) for heparin neutralization in coronary artery surgery.
Heres E K; Horrow J C; Gravlee G P; Tardiff B E; Luber J AUTHOR:

Jr; Schneider J; Barragry T; Broughton R
Allegheny General Hospital, Pittsburgh, Pennsylvania, USA.
ANESTHESIA AND ANALGESIA, (2001 Dec) 93 (6) 1446-52, table CORPORATE SOURCE: SOURCE:

of contents.

Journal code: 1310650. ISSN: 0003-2999.

PUB. COUNTRY:

JOURNAL COURSE STATES OF S

LANGUAGE: English

FILE SEGMENT: ENTRY MONTH: Abridged Index Medicus Journals; Priority Journals

ENTRY DATE:

L14 ANSWER 2 OF 14 ACCESSION NUMBER: MEDLINE

2001321665 MEDLINE

21201502 PubMed ID: 11304642 Analysis of the hemodynamic effects of heparinase I and DOCUMENT NUMBER:

TITLE:

protamine sulfate in the systemic and hindlimb vascular beds of the male rat. Jahr J S; Stuart J S

AUTHOR:

Department of Anesthesiology, University of California Davis Medical Center, Sacramento, CA 95817, USA. AMERICAN JOURNAL OF THERAPEUTICS, (2000 Nov) 7 (6) 353-7. Journal code: DB7; 9441347. ISSN: 1075-2765. CORPORATE SOURCE:

SOURCE:

PUB. COUNTRY: United States

LANGUAGE:

Journal, Article; (JOURNAL ARTICLE)
English
Priority Journals
200106 FILE SEGMENT: ENTRY MONTH:

Entered STN: 20010611 ENTRY DATE:

AY MONTH: 200106

INT DATE: Entered STN: 20010611

Last Updated on STN: 20010611

Entered Medline: 20010607

The effects of heparinase I and protamine sulfate on the mean arterial pressure and hindlimb perfusion pressure in the male rat were studied. With institutional approval, 21 male Sprague-Dawley rats were anesthetized, and the carotid artery and abdominal aorta were cannulated by cutdown. To isolate the hindlimb, a semi-closed peristaltic perfusion circuit was used. Heparinase I or protamine sulfate was injected into the hindlimb vascular bed, and changes in mean arterial pressure and hindlimb perfusion pressure were recorded. Analysis of variance with a post hoc Scheffe's test was used for statistical analysis, and a P value less than .05 was considered significant. Increasing doses of heparinase I caused a small but significant decrease in mean arterial pressure only at the two highest doses. At all doses, hindlimb perfusion pressure was significantly less than the baseline value and than the value with saline administration at 1 minute. At the clinically applicable doses of heparinase I (0.625 and 1.25 IU/kg), the decrease in hindlimb perfusion pressure was less than 7. At the next two higher doses, the change was less than 15k. The vehicle of heparinase caused a significant decrease in mean arterial pressure (from -15k to -30k) and hindlimb perfusion pressure (from -10k to -20k). Increasing doses of protamine sulfate caused an increase in hindlimb perfusion pressure from baseline, including a 58k change with the 10-mg/kg dose. There was a transient decrease in mean arterial pressure, which peaked 4 to 5 minutes after injection, to a 21k decrease from baseline with the 5- and 10-mg/kg doses.

Heparinase I caused vasodilation in the hindlimb and decreased mean arterial pressure only at supraclinical doses. Protamine sulfate caused a significant dose-dependent increase in hindlimb vascular resistance and a transitory decrease in mean arterial pressure.

. . . doses. At all doses, hindlimb perfusion pressure was significantly less than the baseline value and than the value with saline administration at 1 minute. At the clinically applicable doses of heparinase I (0.625 and 1.25 IU/kg), the decrease in hindlimb perfusion pressure was less than 7. At the next two higher. . .

ANSWER 3 OF 14 MEDLINE

AUTHOR:

2001141460 MEDLINE ACCESSION NUMBER: DOCUMENT NUMBER:

21091197 PubMed ID: 11159222 Pentalyte does not decrease heparinoid release but does TITLE:

Pentalyte does not decrease heparinoid release but does decrease circulating thrombotic mediator activity associated with aortic occlusion-reperfusion in rabbits. Nielsen V G; Armstead V E; Geary B T; Opentanova I L Department of Anesthesiology, Division of Cardiothoracic Anesthesia, The University of Alabama at Birmingham, Birmingham, Alabama 35249, USA.. vance.nielsen@ccc.uab.edu KOB-GM00686 (NIGMS)

CORPORATE SOURCE:

CONTRACT NUMBER:

ROB-CHUUUBS (NIGMS) ANESTHESIA AND ANALGESIA, (2001 Feb) 92 (2) 314-9. Journal code: 4R8; 1310650. ISSN: 0003-2999. SOURCE:

PUB. COUNTRY: United States

Journal; Article; (JOURNAL ARTICLE) LANGUAGE: English

FILE SEGMENT: ENTRY MONTH:

Abridged Index Medicus Journals; Priority Journals 200103

Entered STN: 20010404 ENTRY DATE:

Last Updated on STN: 20010404 Entered Medline: 20010308

Last Updated on STN: 20010404
Entered Medline: 20010308

Hemorrhage and thrombosis are associated with major vascular and trauma surgery. Release of heparinoids and thrombotic mediators may contribute to these complications and have been described in rabbits after aortic occlusion-reperfusion. We hypothesized that the resuscitative fluid used could reduce heparinoid and thrombotic mediator release after aortic occlusion-reperfusion in rabbits as assessed by thromboelastographic variables (R, reaction time; alpha, angle; and G, a measure of clot strength). Anesthetized rabbits were administered lactated Ringer's solution (n = 8) or Pentalyte (n = 8) at reperfusion after 30 min of ischemia. Blood was obtained before ischemia and after 30 min of reperfusion for thromboelastography under four conditions: 1) unmodified sample, 2) platelet inhibition, 3) heparinase, and 4) platelet inhibition and heparinase. During reperfusion, unmodified samples demonstrated a significant increase in R and decrease in alpha and G that was not affected by Pentalyte. In the presence of heparinase, no significant fluid-specific thromboelastographic differences were noted. However, thrombotic mediator release (discerned by a decrease in R and an increase in alpha) during reperfusion in samples with platelet inhibition and heparinase was significantly attenuated by Pentalyte. Pentalyte administration does not decrease heparinoid release but does decrease thrombotic mediator release after aortic occlusion-reperfusion.

. release (discerned by a decrease in R and an increase in alpha) during reperfusion in samples with platelet inhibition and heparinase was significantly attenuated by Pentalyte. Pentalyte Administration does not decrease heparinoid release but does decrease thrombotic mediator release after aortic occlusion-reperfusion.

Answer 4 OF 14 MEDLINE

L14 ANSWER 4 OF 14 ACCESSION NUMBER:

2001034941 MEDLINE DOCUMENT NUMBER:

20534203 PubMed ID: 11083232 Influence of the endothelial glycocalyx on cerebral blood TITLE:

flow in mice. AUTHOR:

J; Sperandio M; Pries A R; Linderkamp O; Gaehtgens P; Kuschinsky W Department of Physiology and Pathophysiology, University of

CORPORATE SOURCE:

SOURCE:

Heidelberg, Germany. JOURNAL OF CEREBRAL BLOOD FLOW AND METABOLISM, (2000 Nov) 20 (11) 1571-8.

Journal code: HNL. ISSN: 0271-678X. United States PUB. COUNTRY: Journal; Article; (JOURNAL ARTICLE)

English Priority Journals

LANGUAGE: FILE SEGMENT: ENTRY MONTH: 200011

MONTH: 200011

(DATE: Entered STN: 20010322

Last Updated on STN: 20010322

Entered Medline: 20001130

The endothelial surface layer (glycocalyx) of cerebral capillaries may increase resistance to blood flow. This hypothesis was investigated in mice by intravenous administration of heparinase (2500 mice by intravenous administration of negatinase (2500 IV/kg body weight in saline), which cleaves proteoglycan junctions of the glycocalyx. Morphology was investigated by transmission electron microscopy. Cerebral perfusion velocity was recorded before and during heparinase or saline treatment using laser-Doppler flowmetry. In addition, cerebral blood flow (CBF) was measured 10 minutes after heparinase or cerebral blood flow (CBF) was measured 10 minutes after heparinase or saline treatment using the iodo[14C] antipyrine method. Laser-Doppler flowmetry and CBF measurements were performed during normocapnia and severe hypercapnia (PCO2: 120 mm Hg). After heparinase, morphology showed a reduced thickness of the glycocalyx in cortical microvessels by 43% (P < 0.05) compared with saline-treated controls. Under normocapnic conditions, a 15% (P < 0.05) transient increase of cerebral flow velocity occurred 2.5 to 5 minutes after heparinase injection. Laser-Doppler flow and CBF returned to control values ten minutes after the injection. However, returned to control values ten minutes after the injection. However, during severe hypercapnia, heparinase treatment resulted in a persisting increase in laser-Doppler flow (64, P < 0.05) and CBF (304, P < 0.05). These observations indicate the existence of a flow resistance in cerebral capillaries exerted by the glycocalyx. The transient nature of the CBF increase during normocapnia may be explained by a vascular compensation that is exhausted during severe hypercapnia. . . . surface layer (glycocalyx) of cerebral capillaries may increase resistance to blood flow. This hypothesis was investigated in mice by intravenous administration of heparinase (2500 IU/kg body weight in saline), which cleaves proteoglycan junctions of the glycocalyx. Morphology was investigated by transmission electron microscopy. . .

ANSWER 5 OF 14 MEDLINE

ACCESSION NUMBER: DOCUMENT NUMBER:

microscopy...

2000286215 MEDLINE

200266215 PubMed ID: 10825314
Rapid evaluation of coagulopathies after cardiopulmonary bypass in children using modified thromboelastography. TITLE:

CORPORATE SOURCE:

Miller B E; Guzzetta N A; Tosone S R; Levy J H Department of Anesthesiology, Emory University School of Medicine, Atlanta, Georgia 30322, USA. ANESTHESIA AND ANALGESIA, (2000 Jun) 90 (6) 1324-30.

SOURCE:

Journal code: 4R8; 1310650. ISSN: 0003-2999.

United States (CLINICAL TRIAL) PUB. COUNTRY:

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: English

Abridged Index Medicus Journals; Priority Journals 200006

FILE SEGMENT: ENTRY MONTH:

Entered STN: 20000629 ENTRY DATE:

Last Updated on STN: 20000629 Entered Medline: 20000616

Entered Meddine: 20000616

Complex coagulopathies follow cardiopulmonary bypass (CPB) in children. However, objective laboratory data that can be acquired rapidly to guide their management are lacking. Because thromboelastography has proven useful in this regard, we evaluated the use of celite or tissue factor (TF) activation and heparinase modification of blood samples to allow rapid determination of thromboelastogram data in children younger than 2 yr undergoing CPB. Celite or TF activation shortened the initiation of clotting and, thus, the time required for the important thromboelastogram alpha and maximum amplitude values to begin evolving. Although thromboelastogram alpha and maximum amplitude values were increased with these activators, correlations persisted between platelet count or fibrinogen level and each of these values. The additional use of heparinase allowed thromboelastograms to be obtained during CPB with values not different from those obtained without heparinase after protamine administration. Therefore, celite- or TF-activated, heparinase-modified thromboelastograms begun during CPB allow objective data to be available by the conclusion of protamine administration to help restore hemostasis after CPB in children. Thromboelastography identified transient fibrinolysis during CPB in some children that resolved by the conclusion of protamine administration. Future investigations of the effectiveness of modified thromboelastography-guided coagulopathy management after CPB in children represended lumilistration. Thromboelastography assessing the Future investigations of the effectiveness of modified thromboelastography-guided coagulopathy management after CPB in children are needed. Implications: Thromboelastography is useful in assessing the coagulopathies that follow cardiopulmonary bypass in children. Modifying blood samples with celite or tissue factor and heparinase allows thromboelastography begun before the termination of cardiopulmonary bypass to become a rapid point-of-care monitor to provide objective data for guiding blood component therapy to manage these coagulopathies.

. . The additional use of heparinase allowed thromboelastograms to be obtained during CPB with values not different from those obtained without heparinase after protamine administration. Therefore, celite- or TF-activated, heparinase-modified thromboelastograms begun during CPB allow objective data to be available by the conclusion of protamine administration to help restore hemostasis.

protamine administration to help restore hemostasis.

ANSWER 6 OF 14 MEDLINE

ACCESSION NUMBER: DOCUMENT NUMBER: 1999384172 MEDLINE 99384172 PubMed ID: 10454476

99384172 PubMed ID: 10454476
Ex vivo reversal of heparin-mediated cardioprotection by heparinase after ischemia and reperfusion.
Kilgore K S; Tanhehco E J; Naylor K B; Lucchesi B R
Department of Pharmacology, University of Michigan Medical School, Ann Arbor, Michigan, USA.
JOURNAL OF PHARMACOLOGY AND EXPERIMENTAL THERAPEUTICS, (1999 Sep) 290 (3) 1041-7.
Journal code: JF3; 0376362. ISSN: 0022-3565.
United States
Journal: Article: (JOURNAL ARTICLE) CORPORATE SOURCE:

SOURCE:

PUB. COUNTRY:

Journal; Article; (JOURNAL ARTICLE)

LANGUAGE: Priority Journals FILE SEGMENT:

AUTHOR

ENTRY MONTH: ENTRY DATE:

199909 Entered STN: 19991005

Last Updated on STN: 19991005 Entered Medline: 19990921

Last Updated on STN: 19991005
Entered Medline: 19990921

Glycosaminoglycans, including heparin, have been demonstrated both in vitro and in vivo to protect the ischemic myocardium against reperfusion injury. In the present study, we sought to determine whether the cardioprotective effects of heparin administration could be reversed by the heparin-degrading enzyme heparinase. New Zealand white rabbits were pretreated with heparin (300 U/kg i.v.) or vehicle (saline). Two hours after treatment, hearts were removed, perfused on a Langendorff apparatus, and subjected to 25 min of global ischemia, followed by 45 min of reperfusion. Hemodynamic variables were obtained before ischemia (baseline) and every 10 min throughout the reperfusion period. Compared with vehicle-treated rabbits, the left ventricular end-diastolic and left ventricular developed pressures were improved significantly (p<.05) in the heparin-treated group. Ex vivo administration of heparinase (5 U/ml) immediately before the onset of global ischemia was associated with a reversal of the heparin-mediated cardioprotection. The uptake of a radiolabeled antibody to the intracellular protein myosin and creatine kinase release were used to determine membrane integrity and discriminate between viable and nonviable myocardial tissue. The uptake of radiolabeled antimyosin antibody and release of creatine kinase after reperfusion were increased in heparin-pretreated hearts exposed to heparinase, indicating a loss of membrane integrity and increased myocyte injury. These results demonstrate that neutralization of heparin by heparinase promotes increased myocardial injury after reperfusion of the ischemic myocardium.

. . . the ischemic myocardium against reperfusion injury. In the present study, we sought to determine whether the cardioprotective-effects of heparin administration could be reversed by the

the ischemic myocardium against reperfusion injury. In the present study, we sought to determine whether the cardioprotective-effects of heparin administration could be reversed by the heparin-degrading enzyme heparinase. New Zealand white rabbits were pretreated with heparin (300 U/kg i.v.) or vehicle (saline). Two hours after treatment, hearts were. . . the left ventricular end-diastolic and left ventricular developed pressures were improved significantly (p <.05) in the heparin-treated group. Ex vivo administration of heparinase (5 U/ml) immediately before the onset of global ischemia was associated with a reversal of the heparin-mediated cardioprotection. The uptake.

L14 ANSWER 7 OF 14 ACCESSION NUMBER:

DOCUMENT NUMBER: TITLE:

MEDLINE
198300542 MEDLINE
98300542 PubMed ID: 9636913
Thromboelastography with heparinase in orthotopic liver transplantation.
Pivalizza E G; Abramson D C; King F S Jr
Department of Anesthesiology, University of Texas Health Science Center, Houston 77030, USA. CORPORATE SOURCE:

JOURNAL OF CARDIOTHORACIC AND VASCULAR ANESTHESIA, (1998 Jun) 12 (3) 305-8.
Journal code: A6I; 9110208. ISSN: 1053-0770. SOURCE : United States PUB. COUNTRY: Journal; Article; (JOURNAL ARTICLE) LANGUAGE: FILE SEGMENT: English Priority Journals ENTRY MONTH: ENTRY DATE: 199809 Y DATE: 19980917
Y DATE: Entered STN: 19980917
Last Updated on STN: 19980917
Entered Medline: 19980909
OBJECTIVE: To investigate the role of heparin in the postreperfusion OBJECTIVE: To investigate the role of heparin in the postreperfusion coagulopathy during liver transplantation with heparinase-guided thromboelastography. DESIGN: A prospective, interventional study. SETTING: A university-affiliated hospital. PARTICIPANTS: Twenty-six patients undergoing orthotopic liver transplantation (OLT). INTERVENTIONS: Blood drawn at five intervals for thromboelastography assessment with native (12 patients) or celite blood (14 patients) compared with simultaneous thromboelastography traces with added heparinase. MAIN RESULTS: In the native samples, the prolonged R (reaction) and K (coagulation) time and decreased alpha angle were corrected in heparinase thromboelastograph traces immediately before reperfusion and 10 minutes postreperfusion. In the celite-accelerated samples, the heparinase traces showed correction of the R and K times and alpha angle only at the 10-minute postreperfusion stage. In seven patients who had thromboelastography performed after protamine administration, there were no differences between celite and heparinase-celite traces. CONCLUSIONS:
Heparinase-treated thromboelastography offered compelling evidence for the presence of heparin-like activity after liver graft reperfusion. The objective evidence provided by this modification of thromboelastography-guided protamine administration and was useful in identifying one of the many potential causes of postreperfusion bleeding in patients undergoing many potential causes of postreperfusion bleeding in patients undergoing out.

. . K times and alpha angle only at the 10-minute postreperfusion stage. In seven patients who had thromboelastography performed after protamine administration, there were no differences between celite and heparinase-celite traces. CONCLUSIONS:
Heparinase-treated thromboelastography offered compelling evidence for the presence of heparin-like activity after liver graft reperfusion. The objective evidence. . . L14 ANSWER 8 OF 14 ACCESSION NUMBER: MEDLINE 97097806 97097806 MEDLINE DOCUMENT NUMBER: PubMed ID: 8942349 Heparinase-guided thrombelastography in an anticoagulated Heparinase-guided thrombelastography in an anticoagulated parturient.

Abramson D C; Abouleish E I; Pivalizza E G; Luehr S L; Myers T; Phillips M Department of Anesthesiology, University of Texas Medical School at Houston 77030, USA.

BRITISH JOURNAL OF ANAESTHESIA, (1996 Oct) 77 (4) 556-8.

JOURNAL Oct. AUO; 0372541. ISSN: 0007-0912.

ENGLAND: United Kingdom

LOWERS AND ARTICLE N. AUTHOR: CORPORATE SOURCE: SOURCE: PUB. COUNTRY: Journal; Article; (JOURNAL ARTICLE) LANGUAGE: English FILE SEGMENT: Priority Journals Entered STN: 19970128 Entered STN: 19970128

Last Updated on STN: 19980206

Entered Medline: 19961230

We describe the use of heparinase-guided thrombelastography in the assessment of a parturient who had been anticoagulated with heparin for suspected thromboembolic disease. Reversal of the heparin effect in the heparinese-treated sample facilitated administration of protamine and successful subarachnoid analgesia for delivery.

. . of a parturient who had been anticoagulated with heparin for suspected thromboembolic disease. Reversal of the heparin effect in the heparinase-treated sample facilitated administration of protamine and successful subarachnoid analgesia for delivery. ENTRY DATE: AB L14 ANSWER 9 OF 14 ACCESSION NUMBER: MEDLINE 96239965 MEDLINE 96239965 PubMed ID: 8815578 Low molecular weight heparin is responsible for the anti-Xa DOCUMENT NUMBER: activity of Desmin 370. Brieger D; Dawes J Applied Research Group, The Heart Research Institute, AUTHOR CORPORATE SOURCE: Applied Research Group, The Heart Research Institute, Sydney, Australia.

THROMBOSIS AND HAEMOSTASIS, (1996 Peb) 75 (2) 286-91.

JOURNALY: Germany, Federal Republic of Journal, Article; (JOURNAL ARTICLE) SOURCE : PUB. COUNTRY: English LANGUAGE: FILE SEGMENT: ENTRY MONTH: Priority Journals 199610 Entered STN: 19961022 Last Updated on STN: 19980206 ENTRY DATE:

Entered STN: 19961022

Last Updated on STN: 19980206

Entered Medline: 19961010

Dermatan sulphate does not catalyse the inactivation of factor Xa.

However, the low molecular weight (LMW) dermatan sulphate Desmin 370 has been shown to generate circulating anti-Xa activity following administration to humans. Using a single batch of Desmin 370, we measured 3 U/mg of anti-Xa activity by amidolytic assay in vitro. The material responsible for this activity had a lower molecular weight range (6000 and 1800 Da) than Desmin 370 and was more highly sulphated than the bulk of the drug. Heparinase digestion of Desmin 370 eliminated 90% of the in vitro anti-Xa activity without significantly interfering with its ability to potentiate inactivation of thrombin by HCII, suggesting that the anti-Xa activity is not due to dermatan sulphate and is probably heparin. When 1251-labelled Desmin 370 together with 40 mg/kg carrier drug was administered intravenously to a rabbit, anti-Xa activity was readily detectable in the plasma for up to 10 h and had a longer half-life than the sulphated radiolabel. Most of this anticoagulant activity was recovered from the plasma by Polybrene affinity chromatography and was probably a sulphated glycosaminoglycan. Administration of the heparinase-digested drug to a rabbit resulted in 70% less anti-Xa activity than the undigested drug. We conclude that Desmin 370 contains detectable quantities of biologically active low molecular weight heparin, which is responsible for persistent anti-Xa activity following intravenous administration. administration. Most of this anticoagulant activity was recovered from the plasma by Polybrene affinity chromatography and was probably a sulphated

glycosaminoglycan. Administration of the maparinase -digested drug to a rabbit resulted in 70% less anti-Xa activity than the undigested drug. We conclude that Desmin 370 contains. . . ANSWER 10 OF 14 MEDLINE ACCESSION NUMBER: DOCUMENT NUMBER: 87100328 MEDLINE 87100328 PubMed ID: 3801083 Bffect of very low molecular weight heparin-derived oligosaccharides on lipoprotein lipase release in rabbits. Merchant Z M; Erbe E E; Eddy W P; Patel D; Linhardt R J ATHEROSCLEROSIS, (1986 Nov) 62 (2) 151-8.
JOURNAL code: 95X; 0242543. ISSN: 0021-9150. TITLE: AUTHOR PUB. COUNTRY: Netherlands

Journal; Article; (JOURNAL ARTICLE)

English Priority Journals LANGUAGE . ENTRY MONTH: 198701 ENTRY DATE: Entered STN: 19900302 Last Updated on STN: 19980206

Entered STN: 19900302

Last Updated on STN: 19980206

Entered Medline: 19870130

Oligosaccharide fragments of heparin were prepared using flavobacterial heparinase. Following sizing, these oligosaccharide fractions were administered (i.v.) to rabbits and were examined for their ability to release lipoprotein lipase. The decasaccharides (dp = 10, Mr avg = 2,800) were the smallest oligosaccharides which resulted in substantial lipase release. The plasma lipase levels obtained with decasaccharides were comparable to low molecular weight heparin and one-third those obtained when heparin was administered at an equivalent dose. The peak plasma lipase concentration was observed 10 min following heparinization and fell off rapidly over the 60-min time course. The lipase release activity paralleled the in vivo pharmacokinetics of the heparin and decasaccharide sample as determined by monitoring their anti-Factor Xa activity. No activation of purified bovine milk lipoprotein lipase or plasma lipase was detectable at the concentrations studied, indicating that the increase in circulating lipolytic activity was due entirely to release. Lipoprotein lipase accounted for a major portion of the released activity with hepatic triglyceride lipase representing the remainder of the lipolytic activity. The sized decasaccharide sample was characterized with regards to its structure and anticoagulant activity. The decasaccharides exhibited reduced anticoagulant activity possibly making it a better drug candidate in the treatment of atherosclerosis. Oligosaccharide fragments of heparin were prepared using flavobacterial heparinase. Following sizing, these oligosaccharide fractions were administered (i.v.) to rabbits and were examined for their ability to release lipoprotein lipase. The decasaccharides (dp = 10, Mr avg. . .

L14 ANSWER 11 OF 14 CAPLUS COPYRIGHT 2002 ACS ACCESSION NUMBER: 2001:676971 CAPLUS 2001:676971 CAPLUS DOCUMENT NUMBER: 135:221282 Use of heparinase III in cancer treatment and TITLE: INVENTOR(S):

Use of heparinase III in cancer treatment and inhibition of tumor cell growth Dongfang, Liu; Pojasek, Kevin; Shriver, Zachary; Holley, Kristine; El-Shabrawi, Yosuf; Venkataraman, Ganesh; Sasisekharan, Ram Massachusetts Institute of Technology, USA PCT Int. Appl., 94 pp.
CODEN: PIXXD2

PATENT ASSIGNEE(S): SOURCE:

DOCUMENT TYPE: Patent English

LANGUAGE: PAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

WO 2001066772 A2 20010913 WO 2001-US7464 20010308

W: AU, CA, JP
RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL,
PT, SE, TR

PRIORITY APPLN. INFO:

US 2000-187846P P 20000308

AB The invention relates to heparinase III and mutants thereof. Modified forms of heparinase III having reduced enzymic activity which are useful for a variety of purposes, including sequencing of heparin-like glycosaminoglycans (HLGAGs), removing active heparan sulfate from a soln., inhibition of angiogenesis, etc. have been discovered according to the invention. The invention in other aspects relates to methods of treating cancer and inhibiting tumor cell growth and/or metastasis using heparinase III, or products produced by enzymic cleavage by heparinase III of HLGAGS.

IT Antitumor agents

(heparinase II administration with; use of heparinase III in cancer treatment and inhibition of tumor cell growth)

ANSWER 12 OF 14 CAPLUS COPYRIGHT 2002 ACS SSION NUMBER: 1983:27563 CAPLUS

DOCUMENT NUMBER:

SOURCE:

AUTHOR (S):

CORPORATE SOURCE:

98:27563
In vivo activity of microbial heparinase
Langer, R.; Linhardt, R. J.; Larsen, A. K.; Cooney, C.
L.; Tapper, D.; Klein, M.
Dep. Nutr. Food Sci., MIT, Cambridge, MA, USA
Trans. - Am. Soc. Artif. Intern. Organs (1982), 28,
387-90
COREN. TATORY AND TATORY CONTROL TATORY AND TATORY

CODEN: TAIOAL; ISSN: 0066-0078

DOCUMENT TYPE:

LANGUAGE: English

NAGE: English
In rabbits, i.v. administration of heparinase
[9025-39-2] (0.5 mg) either 1 min before or 1 min after heparin
[9005-49-6] administration substantially decreased the activated partial
thromboplastin time (aPTT). Blood from heparinized dogs was passed
through a filter of immobilized heparinase. Within 2 min or 1 pass
through the filter nearly all the anticoagulant activity of heparin (aPTT)
was destroyed. Antibodies to heparin were not detected in the blood of
the dogs 2 mo after the expt.
In rabbits, i.v. administration of heparinase
[9025-39-2] (0.5 mg) either 1 min before or 1 min after heparin
[9005-49-6] administration substantially decreased the activated partial
thromboplastin time (aPTT). Blood from heparinized dogs was passed
through a filter of immobilized heparinase. Within 2 min or 1 pass
through the filter nearly all the anticoagulant activity of heparin (aPTT)
was destroyed. Antibodies to heparin were not detected in the blood of
the dogs 2 mo after the expt.

ACCESSION NUMBER: 97017864 EMBASE 1997017864 DOCUMENT NUMBER: 1997017864
Do heparinase thrombelastographs predict postoperative bleeding?.
Mashburn P.; Ecklund J.; Riley J.
J. Ecklund, MUSC-ECT, 101 Doughty St., Charleston, SC 29401, United States
Journal of Extra-Corporeal Technology, (1996) 28/4 (185-190). AUTTHOR CORPORATE SOURCE: SOURCE: Refs: 10 ISSN: 0022-1058 CODEN: JEXCBD United States COUNTRY: Journal; Article 025 Hematology DOCUMENT TYPE: FILE SEGMENT: 025 LANGUAGE: SUMMARY LANGUAGE: English English Postoperative hemorrhage is a major cause of morbidity and mortality in patients who undergo cardiopulmonary bypass (CPB). The thrombelastograph (TEG) is a viscoelastic whole blood test that measures clot dynamics from Postoperative hemorrhage is a major cause of morbidity and mortality in patients who undergo cardiopulmonary bypass (CPB). The thrombelastograph (TEG) is a viscoelastic whole blood test that measures clot dynamics from clot formation through clot lysis. Previous studies have shown that post-bypass TEGs are accurate predictors of postoperative bleeding. TEGs from heparinized blood reversed with heparinase may be employed during CPB to evaluate coagulation. CPB heparinase TEGs may allow for earlier recognition of patients who may bleed after bypass. Earlier TEG analysis would allow targeting of specific therapies to begin before the patient bleeds excessively. Fifty-four heparinase TEGs during warming and fifty-four native TEGs post-protamine administration were collected. Parameters evaluated were R, K, alpha angle, MA, MA6, coagulation index, activated clotting time, hematocrit, prothrombin time, partial thromboplastin time, thrombin time, fibrinogen concentration, platelet count, blood loss during and after CPB, and blood and blood product administration. Coagulation indexes for CPB heparinase TEGs that were fibrinolytic were 87% accurate in predicting patients with excessive intraoperative blood loss, but were not predictive of blood product administration. The sensitivity was 12.5% and the specificity was 100% in predicting excessive intraoperative bleeding. Post-protamine coagulation index inversely correlated with intraoperative red blood cell administration (r=0.403, p<0.05), but was not predictive. Patients with fibrinolytic TEGs required blood products to compensate for expected blood loss associated with the fibrinolytic state. Simultaneous routine coagulation tests did not correlate significantly with blood loss or blood product administration, nor were they predictive. The findings of this study suggest that the presence of fibrinolysis in either a heparinase TEG on bypass or a post-protamine TEG is the most important predictor of blood and blood product administration. But, since only 20% of the pat predicting patients with excessive. BIOSIS COPYRIGHT 2002 BIOLOGICAL ABSTRACTS INC. 2002:151746 BIOSIS PREV200200151746 L14 ANSWER 14 OF 14 ACCESSION NUMBER: DOCUMENT NUMBER: PREVZ00200151746
Neutralization of the anticoagulant and hemorrhagic effects
of an ultra-low molecular weight heparin (OP2000) by
heparinase-I: Potential clinical implications.
Shadid, H. (1); Daud, A. N.; Iqbal, O.; Ahmad, S.;
Hoppensteadt, D. A.; Demir, M.; Walenga, J. M.; Ward, D.
P.; Fareed, J. (1) TITLE: AUTHOR (S): P.; Fareed, J. (1) (1) Pharmacology, Loyola University of Chicago, Maywood, IL CORPORATE SOURCE: SOURCE: (November 16, 2001) Vol. 98, No. 11 Part 2, pp. 99b. Blood, http://www.bloodjournal.org/. print.
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An ultra-low molecular weight heparin, OP2000, from porcine mucosal heparin has shown better efficacy than unfractionated heparin (UFH) in the management of unstable angina (Clin. Cardiol. 22:213-7, 1999). This drug is currently being developed for the treatment of inflammatory bowel disease, particularly in ulcerative colitis. The treatment dosage for both of these indications is 150 mg/day. While the relative bleeding effects of this agent are lower than UFH because of the significant anti-IIa component and release of tissue factor pathway inhibitor (TPPI), the hemorrhagic potential exists. Protamine sulfate produces a partial neutralization of this agent. In this study, heparinase-I was used to neutralize the anticoagulant and bleeding effect of this agent in experimental and animal models. OP2000 was supplemented to whole blood in a concentration range of 0-100 mcg/ml. Activated clotting time (ACT) and thromboelastography (TEG) tests were performed to study the anticoagulant effects of OP2000. A concentration-dependent increase in the ACT and modification of the TEG parameters were noted. The neutralizing effect of heparinase-I was studied in the concentration range of 0-05-5.0 U/ml. It was observed that heparinase-I at concentration of 2 U/ml produced a complete neutralization of the anticoagulant effect of OP2000 was noted in the activated partial thromboplastin time (APTT), Heptest, and anti-IIa activities. The molecular profiling of OP2000 by GPC-MPLC prior to and after the heparinase digestion revealed marked differences. The average molecular weight was reduced to di- and tri-saccharide components. At a dosage of 10 mg/kg i.v., OP2000 produced prolongation of the bleeding time in animal models (rats, rabbits, primates). Administration of heparinase-I at 0.25 U/kg markedly reversed the bleeding effects of these animal models. In contrast to the heparinase-I studies, protamine sulfate exhibited partial neutralization of OP2000 in these models. These

effective antagonist for the neutralization of this compound.
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3125 S BENNET D?/AU OR CAUCHON E?/AU OR FINK D?/AU OR GROUIX B?/AU O

43 S L1 AND HEPARIN OR HEPARINSE

21 DUP REM L2 (22 DUPLICATES REMOVED)

7 S L3 (P) ADMINIST?

0 S L4 (P) (IV OR INTRAVASCULAR?)

59 S L1 AND (HEPARINASE OR HEPARIN)

29 DUP REM L6 (30 DUPLICATES REMOVED)

7 S L7 (P) ADMINIST?

0 S L8 NOT L4

3653 S (HEPARINASE OR HEPARINASE?)

41 S L10 (10N) (ADMINIST?)

0 S L11 (P) (IV OR INTRAVASCUL?)

18 DUP REM L11 (23 DUPLICATES REMOVED)

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